Amphiphilic protein micelles for targeted in vivo imaging

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Abstract

A variety of polymeric nanoparticles have been developed for bioimaging applications. This study reports on the use of a 50 nm recombinant protein nanoparticle with a multivalent surface as a vehicle for functionalization with a model imaging agent. Multiple fluorescent probes were covalently conjugated to surface amines of crosslinked amphiphilic elastin-mimetic protein micelles using N-hydroxysuccinimide ester chemistry. In vivo fluorescence imaging confirmed that protein micelles selectively accumulated at sites of angioplasty induced vessel wall injury, presumably via an enhanced permeability and retention effect. This investigation demonstrates the potential of amphiphilic protein micelles to be used as a vehicle for selective imaging of sites associated with a disrupted or leaky endothelium.

1. Introduction

Fluorescent nanoparticles have been used to localize and assess the extent of a variety of pathological processes [1–4]. In this regard, both encapsulation and covalent conjugation of fluorophores have been used to generate fluorescent nanoparticles [5–7]. Even without surface targeting groups, nanoparticles selectively accumulate at sites of increased vascular permeability, often associated with tumor microcirculation or inflammatory processes via an enhanced permeability and retention (EPR) effect [2,8–10].

Recent strategies to create probes for biomedical imaging have included organic and inorganic nanoparticles, quantum dots (QD), liposomes, proteins and viral particles [2,3,11–16]. Although each approach has unique advantages, none has proved ideal, owing to concerns related to toxicity, biostability and bioavailability. Among these probes, QD are the most interesting class of fluorescent probes for bioimaging, because of their brightness, photostability and narrow and tunable emission spectrum. In addition to QD, magnetic nanoparticles such as iron oxides have been used as contrast enhancing agents for magnetic resonance imaging (MRI) and coupled with fluorescent probes for both MRI and fluorescence imaging. However, under physiological conditions, QD and magnetic nanoparticles are not soluble and tend to aggregate. In addition, Cd, Te, Se or Pb found in QD are cytotoxic [17]. Protein-based nanoparticles may have several advantages over other imaging agents, including limited toxicity and enhanced biocompatibility, particularly for elastin-based micro- or nanoparticles [18–20]. Significantly, such particles display inherent flexibility for functionalization with bioactive molecules either chemically or genetically [21–24]. For example, Simnick et al. [24] recently reported the generation of elastin-like diblock copolymers, engineered with an RGDS peptide sequence at the N terminus and one cysteine residue at the C terminus, which can be chemically conjugated to a fluorescent probe containing maleimide. The present authors’ group recently developed recombinant amphiphilic diblock polypeptides (ADP) based on elastin-mimetic sequences consisting of an N-terminal hydrophilic block and a C-terminal hydrophobic block containing glutamic acid and tyrosine residues, respectively. It was demonstrated that these polypeptides formed 50-nm-diameter thermally responsive micellar nanoparticles that exhibited a spherical core–shell structure. By introducing multiple cysteine residues between the amphiphilic blocks, it was possible to obtain stable protein micelles through disulfide bond formation at the core–shell interface [25,26].

This study describes the surface functionalization of elastin-mimetic protein (EMP) micelles via chemical conjugation and the potential to use these particles for in vivo bioimaging. Each diblock polypeptide possesses a single free amine at the N terminus, which is displayed on the surface of the protein micelle owing to self-assembly of C-terminal hydrophobic blocks that occurs above their inverse transition temperature. Thus, it was anticipated that functionalizing EMP micelles with fluorescent dyes via an N-hydroxysuccinimide (NHS) ester linker would generate protein nanoparticles with enhanced fluorescent intensity.
Interventional vascular procedures such as balloon angioplasty and stent implantation lead to endothelial injury, which may contribute to subsequent restenosis and arterial occlusion. An effective non-invasive modality to image the extent of endothelial denudation after procedures does not exist. To evaluate the utility of fluorescently labeled protein micelles, the present authors studied their use in detecting discrete sites of vascular wall injury. Balloon injury results in the loss of the endothelium, with significant local enhancement of vessel wall permeability. The EPR effect after angioplasty has been demonstrated using Evans Blue staining, which associates with serum albumin, as well as with hydrophobic drug-conjugated nanoparticles [2,27,28]. It was hypothesized that accumulation of fluorescent EMP micelles could be used as a non-invasive marker for EPR, which characterizes vascular wall injury. In this investigation, EMP micelles were evaluated in a rat aortic balloon injury model with micelle uptake visualized by an in vivo fluorescent imaging system. To the authors’ knowledge, this is the first report in which protein nanoparticles have been used to image sites of vessel wall injury via an EPR effect.

2. Materials and methods

2.1. Materials

All chemical reagents were purchased from Fisher Scientific, Inc. (Pittsburgh, PA) unless otherwise noted. Texas Red d6-X succinimidyl ester or fluorescein succinimidyl ester were purchased from invitrogen (Eugene, Oregon), TALON metal affinity resin was purchased from BD Biosciences, Inc. Methods used to produce the diblock genes encoding ADP derived from elastin-mimetic sequences, (VPVGV)(VPVGE)(VPVGV)(VPVGE)(VPVGV)10 for the hydrophilic N-block and (IPGVG)2(IPGVG(IPGVG)15 for the hydrophobic C-block, have been described previously [24]. Expression of the diblock synthetic genes with N-terminal his-tag sequence in Escherichia coli expression strain, BL21(DE3), afforded recombinant ADP from the expression strain, BL21(DE3), afforded recombinant ADP (1 mg mL−1 culture) when 1 mM isopropyl β-D-1-thiogalactopyranoside was added. Purification was performed by immobilized metal affinity chromatography from the cell lysate. Aqueous solutions of ADP were prepared from lyophilized specimens of the purified protein polymer, dissolved at the appropriate concentration in distilled, deionized water or phosphate buffered saline (PBS) at 4 °C. The purified protein polymer was characterized by sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE), matrix-assisted laser desorption/ionization mass, amino acid compositional analysis and 1H nuclear magnetic resonance [25].

2.2. Methods

2.2.1. Preparation and characterization of elastin-mimetic micelles

Stock solutions of ADP were prepared by dissolving the solid protein (1 mg mL−1) in cold water. For preparation of protein micelles, the protein solution was diluted to 0.3 mg mL−1 with PBS and kept on ice for 1 h. The tube containing diluted solution was subsequently moved to a 25 °C water bath and incubated for 30 min. A micelle suspension was stored under constant agitation at room temperature for 2 weeks, and changes in micelle size and size distribution were monitored by dynamic light scattering (DLS) (DynaPro, Protein Solutions).

2.2.2. Labeling protein micelles with fluorescent dye and characterization

N-terminal amines of elastin-mimetic micelles were modified with an aminoreactive fluorescent probe. Solutions of the protein micelles in 1 mL of PBS were prepared at 25 °C. A total of 10 μL of Texas Red d6-X succinimidyl ester or fluorescein succinimidyl ester dissolved in dimethyl sulfoxide at a concentration of 10 mM was added dropwise to ADP solutions. The reaction mixture was stirred at room temperature in the dark overnight and then dialyzed against PBS for 72 h to remove unreacted dyes. For synthesis of dual dye-labeled micelles, fluorescein (FL) succinimidyl ester was added to Texas Red (TR)—micelle conjugate dispersion, and the reaction mixture was stirred at room temperature in the dark overnight. Non-reacted dyes were removed by dialysis.

Fluorescently labeled protein micelles were analyzed by UV spectroscopy, fluorescence spectroscopy, DLS and gel filtration and visualized by illumination on a standard UV light box. UV-vis spectra were recorded on a Cary UV-visible spectrophotometer (Varian Inc.) in a 1 cm quartz cuvette. The degree of labeling was determined from the absorbance of maximum peak of dyes (FL 494 nm and TR 593 nm) using an extinction coefficient (68,000 for FL and 80,000 M−1 cm−1 for TR) relative to concentration of diblock polypeptide. Fluorescence spectra of fluorophore-conjugated micelles were obtained using PCI photon counting spectrofluorometer (ISS Inc.). FL was excited at 494 nm, and the emission spectra were recorded from 500 to 600 nm. TR was excited at 594 nm, and the emission spectra were recorded from 595 to 700 nm. The size and polydispersity of dye–micelle conjugates were analyzed by DLS using a DynaPro (Protein Solutions) at 25 and 37 °C. Sample solutions were filtered through a 0.45 μm nylon syringe filter prior to DLS measurements. DLS measurements were carried out at a fixed scattering angle of 90°. The molecular weight of the protein micelles was estimated from the hydrodynamic radius, using the standard curve of model proteins. The number of single diblock polypeptides associated in one micelle was determined by MWmicelle/MWpolypeptide. Zeta potential was determined from three measurements using DLS (Malvern Zetasizer, zen3600). Gel filtration for size exclusion chromatography (Hi-Prep 16/60, Sephacryl-S100, high resolution) was performed to analyze the homogeneity of protein micelles. The column was pre-equilibrated with PBS buffer, and protein micelles were eluted with a flow rate of 0.6 mL min−1.

2.2.3. Rat balloon injury and Evans Blue staining

Balloon injury was performed to a discrete 10 mm segment of rat thoracic aorta with a 2F Fogarty catheter (10–12-week-old Wistar Rats, 300–400 g; Jackson Laboratory, Bar Harbor, Maine). Induction of anesthesia was performed in a Plexiglas chamber with isoflurane gas anesthetic (5% inhalant to effect). Anesthesia was maintained by isoflurane inhalational anesthetic at 1.5–2% continuous flow. All animals had free access to standard water and chow, and received care and monitoring in compliance with established guidelines (i.e., per the Institutional Animal Care and Use Committee).

Five animals were assessed at each time point after balloon injury (1 h, 7, 14, 21 days). At the time of sacrifice, a jugular venous cut-down was performed and Evans Blue solution (1 mL, 5 mg mL−1) was injected into the venous circulation. One hour after Evans Blue injection, the laparotomy site was reopened, and the thoracic contents were surgically exposed. Saline irrigation was performed via direct access of the left ventricle until liver congestion, color change and intestinal edema were noted. The inferior vena cava was transected to allow for euthanasia via exsanguination. Sharp dissection was used to release the aorta from its surrounding tissues along its length, extending from the aortic arch to the bifurcation. The aortic segment was harvested and analyzed by gross visualization. Injury zones stained dark blue, owing to the uptake of Evans Blue dye. Image J image processing software (NIH, USA) was used to determine the surface area of the blue-stained injury zone. This process was repeated to assess the change in injury zone surface area over time.
2.2.4. Balloon injury of rat aorta and fluorescent imaging

Balloon injury was performed to a discrete 10 mm segment of rat thoracic aorta with a 2F Fogarty catheter (Wistar Rats, 300–350 g; Jackson Laboratory, Bar Harbor, Maine). All animals had free access to standard water and chow, receiving care and monitoring in compliance with guidelines established by the Institutional Animal Care and Use Committee. Immediately following injury, TR–micelles (1 mL, 0.5 mg mL⁻¹ micellar solution in PBS) were injected intravenously. The conjugate was allowed to circulate for 24 h prior to harvest. Harvested samples were imaged enface on an Olympus OV100 fluorescent imaging system. Filters were chosen to encompass the excitation/emissions wavelengths of TR of 595/615 nm, respectively. System parameters (i.e., exposure time, maximum/minimum intensities and focal length) were kept constant for each acquisition, and images were acquired at 6× magnification. With the use of the OV100 software package, heat maps of fluorescent images were generated and exported as .tiff files for further analysis.

A novel image analysis program using MatLab (The Mathworks Inc.) was created to quantify the heat maps. The program allowed for pixel quantification corresponding to each of the eight intensity bars defining the heat map. Each colorimetric bar of the heat map is associated with a fixed mean fluorescent intensity value. The percentage surface area of each of these colors within the injured region was plotted against mean fluorescent intensity to obtain a histogramic representation of the data. Additionally, the eight intensity bars of the heat map were grouped into high (top three bars), medium (middle two bars) and low (bottom three bars) as another means of representing the data with respect to percentage surface area of the stent.

3. Results and discussion

3.1. Conjugation of fluorescent dye to surface of protein micelle

The surface of amphiphilic EMP micelles can be functionalized with a variety of biomolecules, such as fluorescent probes or specific targeting ligands through chemical or recombinant approaches. Amine-reactive molecules were employed for chemical modification, because only a single amino group is available for conjugation in the polypeptide sequence. Formation of spherical micelles with a core–shell structure generated nanoparticles with a multivalent aminated surfaces, owing to the presence of terminal amines in the hydrophilic block. Thus, bifunctional chemical linkers containing NHS ester facilitate the derivatization of surface protein nanoparticles with bioactive molecules.

To examine the feasibility of N-terminal conjugation, fluorescent dyes with an NHS ester moiety, NHS–TR and NHS–FL, were used to modify the surface of amphiphilic EMP micelles (Fig. 1). Briefly, TR succinimidyl ester (NHS–TR) was added to a micellar solution in PBS buffer to generate TR derivatized protein micelles (TR–micelle). The labeling reaction was incubated at room temperature overnight, and unreacted dye was subsequently removed through dialysis against PBS buffer for 72 h (Fig. 2A (left) and B).

3.2. Characterization of fluorescently labeled protein micelle

The protein micelles were characterized by DLS. The number of diblock polypeptides associated in one protein micelle was determined from the molecular weight of the protein micelle [24]. The estimated molecular weight of the spherical protein micelles was 6035 kDa, determined by DLS. Given that the molecular weight of a single diblock polypeptide chain was 56.9 kDa, a single micelle was formed from the assembly of ~106 diblock polypeptide chains with a corresponding number of free amines displayed on the surface. The degree of labeling, as measured by UV-vis spectroscopy, revealed a ratio of dye molecules per mole of diblock polypeptide of 0.5–0.6 consistent with the estimated 50–60 dye molecules per protein micelle, which was also confirmed by fluorescence spectroscopy, using a standard curve based on free dye.

Conjugation of multiple fluorophores to a given protein molecule, such as an antibody, can cause a significant reduction in fluorescence intensity by quenching interactions between closely located dyes. However, large protein particles, such as virus nanoparticles have been labeled with multiple fluorescent dyes, and the dye–virus conjugates generated highly fluorescent particles without fluorescence quenching [15]. Similarly, EMP micelles doped with a number of fluorescent dyes have no observable fluorescence quenching, even after prolonged storage, probably due to the flexible hydrophilic chains and long intermolecular distances between fluorophores.

Labeling of EMP micelles with fluorophores was confirmed by UV-vis spectroscopy and fluorescence spectroscopy (Fig. 2D and E and Fig. S2). Labeling micelles with both dyes were also examined by sequential conjugation of TR–micelles with NHS FL. UV spectroscopy of TR/FL–micelle conjugates exhibited two absorption maxima at 494 and 593 nm, which are characteristic for each fluorophore, respectively (Fig. 2D). Dual dye labeling suggests that EMP micelles could serve as multimodal protein nanoparticles with the incorporation of appropriate imaging agents.

The fluorescently labeled protein micelles were further analyzed by size exclusion chromatography using gel filtration (Fig. 2C). The presence of tyrosine residues in the hydrophobic block facilitated detection of eluted protein micelles. TR- and FL-labeled protein micelles displayed elution times identical to those of non-labeled micelles. As illustrated by DLS and gel filtration profiles, dye conjugation did not dissociate micelle structure or substantially alter size.

After conjugation with a fluorescent probe, the size and stability of fluorescently labeled EMP micelles were monitored by DLS (Fig. 3, Table 1). TR–micelles were stable when stored at room temperature in PBS with constant shaking for 2 weeks. Under physiological conditions, micelle stability remains a critical determinant of the suitability of a given formulation. As a surrogate measure of biostability, DLS was used to assess micelle dimensions in the presence of 10% fetal bovine serum (FBS) at 37 °C. A small increase in polydispersity and micelle size was noted after 1 h incubation.

The zeta potential was measured to study the variation in surface charge. The outer shell of EMP micelles contains glutamic acid residues in the hydrophilic block and an aminated surface. The zeta potential of unlabeled EMP micelles was negative, owing to the highly carboxylated shell (~13 mV). After conjugation of the N-terminal amino group with a fluorescent probe, the zeta potential decreased, owing to a reduction in the number of protonated amino groups on the micelle surface (~18 mV). Therefore, the zeta potential, together with UV and fluorescence spectroscopy, suggested the presence of fluorescent probes on the micelle surface and improved stability of the fluorescently labeled micelles at neutral pH.

3.3. Balloon injury of rat aorta and fluorescent imaging

Interventional vascular procedures are commonly used for the treatment of atherosclerotic occlusive disease. However, such intervention inevitably leads to damage of the vessel wall. Removal of the endothelium by balloon injury results in exposure of collagen to blood and platelet deposition, and may be followed by proliferation and migration of vascular smooth muscle cells. In particular, the risk of restenosis and arterial occlusion remains high after treatment of extensive complex lesions characteristic of lower extremity peripheral arterial disease. One of the interesting biological properties of the vascular endothelium is to mediate...
Fig. 1. Labeling of N-terminal amines of EMP micelles with amine-reactive fluorescent probes, TR and FL NHS esters.

Fig. 2. (A) UV illumination of aqueous dispersion of TR–micelle, TR/FL–micelle and FL–micelle conjugates. (B) SDS-PAGE analysis and UV illumination to detect TR–micelle conjugate. 1, unlabeled protein micelles; 2, mixture of micelles and non-NHS fluorophore (Rhodamine B); 3, TR–micelle conjugate; M, molecular weight standard. (C) Size exclusion chromatographic analysis of unlabeled micelle (dotted), TR–micelle (dashed) and FL–micelle conjugate (solid). (D) UV-visible spectra of unlabeled protein micelle (black), TR–micelle (red), FL–micelle (green) and TR/FL–micelle conjugate (pink). (E) Fluorescence spectra (solid lines) of FL–micelle (green) and TR–micelle (red).
vascular permeability. Previous studies have shown that even marginal disruption of this architecture can significantly enhance vascular wall permeability [27,28]. Given the inevitable endothelial injury during endoluminal vascular interventions, the capacity to detect the site of vessel wall injury may provide a rational approach to monitoring the integrity of endothelium as an initial marker of vascular wall injury. However, there has been no reliable modality for identifying the extent of endothelial denudation after an interventional procedure.

Vascular targeting strategies to detect vessel wall injury include ligand-based targeting and EPR-based targeting. Ligand-based targeting strategy is considered to be effective, but difficult to generate. Among targeting ligands to vascular injury, glycoprotein VI (GPVI) is known as a collagen receptor, which is expressed on platelets and plays a critical role in the activation and aggregation of platelets at the site of injury. A fusion protein of GPVI and a human C-terminal Fc fragment exhibited selective binding to collagen and the site of vessel wall injury in vivo [29,30]. It has been reported recently that the use of FITC–GPVI–Fc (a human Fc fragment) conjugate facilitated the fluorescence imaging of vascular lesions via targeting to collagen exposed to blood flow after endothelial denudation [30]. In contrast, enhanced vascular permeability-based targeting involves the selective uptake of EMP micelles at the site of vessel wall injury.

Prior to in vivo application of the EMP micelles, it was first confirmed that balloon injury of the rat thoracic aorta led to a reproducible model of vessel wall injury [27,28]. Given the inevitable endothelial injury during endoluminal vascular interventions, the capacity to detect the site of vessel wall injury may provide a rational approach to monitoring the integrity of endothelium as an initial marker of vascular wall injury. However, there has been no reliable modality for identifying the extent of endothelial denudation after an interventional procedure.

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Prior to in vivo application of the EMP micelles, it was first confirmed that balloon injury of the rat thoracic aorta led to a reproducible model of vessel wall injury. Endothelial denudation over a 10 mm length with an associated increase in vascular permeability was identified by Evans Blue staining (Fig. 4). The injury zone of the thoracic aorta was stained blue with Evans Blue dye, while the uninjured zone was not stained or stained only pale blue. Serial evaluation of the injury site by Evans Blue analysis revealed that the time to achieve 50% healing ($t_{50}$) was 8.1 days (Fig. S3). Evans Blue staining confirmed the existence of EPR effects consistent with prior reports [2,27].

The biodegradability of EMP micelles is a critical factor for in vivo imaging and its biocompatibility. For in vivo applications, micelles should be stable during circulation to effectively target the site of vessel wall injury but then undergo degradation. Liu et al. determined the in vitro degradation rate after incubation with serum and in vivo degradation rate of 14C-labeled elastin-like polypeptides after intravenous injection, which were 2.49% and 2.46% per day, respectively [31].

The capacity of protein micelles to selectively penetrate the injury site was assessed after IV administration of 0.5 mg of TR-labeled micelles. TR–micelle conjugates were injected in the immediate post-operative period, and imaging was performed after a 24 h circulation time. Significantly, fluorescence imaging demonstrated that micelle uptake correlated with the extent of endothelial denudation and could be used to quantitate regions of increased micelle deposition in the injured vessel wall (Fig. 5A–D). In contrast, micelle uptake was not observed in normal, uninjured aorta (Fig. 5E and F).

**Table 1** Summary of size and polydispersity for unlabeled and TR-labeled protein micelles.

<table>
<thead>
<tr>
<th>Micelle</th>
<th>Size (nm)</th>
<th>PDI a</th>
</tr>
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<tbody>
<tr>
<td>Micelle</td>
<td>49.2</td>
<td>0.17</td>
</tr>
<tr>
<td>Micelle + BSA b</td>
<td>70.0 c</td>
<td>0.12</td>
</tr>
<tr>
<td>Micelle + BSA + TCEP c</td>
<td>172.2 e</td>
<td>0.01</td>
</tr>
<tr>
<td>TR–micelle + FBS d</td>
<td>73.2</td>
<td>0.28</td>
</tr>
<tr>
<td>TR–micelle conjugate ($t$ = 0)</td>
<td>59.0</td>
<td>0.24</td>
</tr>
<tr>
<td>TR–micelle conjugate ($t$ = 2 weeks, PBS, 23°C)</td>
<td>65.3</td>
<td>0.59</td>
</tr>
</tbody>
</table>

a Polydispersity index.

b Bovine serum albumin (3 mg mL$^{-1}$).

c 1 mM Tris(2-carboxyethyl)phosphine.

d 10% FBS.

e Ref. [25].

**Fig. 4.** Balloon injury of rat aorta visualized by Evans Blue staining. Five animals were assessed at each time point after balloon injury (1 h, 7, 14, 21 days). At the time of sacrifice Evans Blue solution (1 mL, 5 mg mL$^{-1}$) was injected into the venous circulation. One hour after dye injection, saline irrigation was performed via direct access of the left ventricle. The aortic segment was harvested, and the injury zone of thoracic aorta was visualized in dark blue color, with non-injured regions indicated by an absence of Evans Blue staining.

**Fig. 3.** DLS measurements of (A) TR–micelle and (B) TR–micelle/10% FBS at 37°C.
4. Conclusions

In conclusion, the present study demonstrates the feasibility of multivalent attachment of fluorophores to amphiphilic EMP micelles with targeting to sites of vessel wall injury. Elastin-mimetic polypeptides have advantages over conventional polymers, including minimal toxicity and good biocompatibility. The surface of EMP micelles was covalently modified with NHS ester modified fluorescent dyes via formation of an amide linkage without substantial fluorescent quenching. The present authors expect that multivalent surface conjugation with a near-IR fluorescence probe or positron emitting agents would further facilitate non-invasive in vivo imaging. Moreover, they believe that multivalent display of targeting ligands including specific cell adhesive peptides or antibodies would further enhance selective particle targeting to distinct cell-surface receptors.

Acknowledgements

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Appendix A. Figures with essential color discrimination

Certain figures in this article, particularly Figs. 1, 2, 4 and 5, are difficult to interpret in black and white. The full color images can be found in the on-line version, at http://dx.doi.org/10.1016/j.actbio.2012.04.011.

Appendix B. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.actbio.2012.04.011.
References


